

Ecology and Conservation of the Endangered Banteng (*Bos javanicus*) in Indonesia Tropical Lowland Forest

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ABSTRACT

Banteng, *Bos javanicus*, as wild cattle is a vital and importance source of germplasm in Indonesia. Various human activities currently threaten their conservation status. Nonetheless, no long-term monitoring programmes are in place for this species. Using distribution point and statistical analysis based on 46,116 camera trap days from December 2015 to January 2017, we aimed to provide habitat preferences, activity patterns and ecological data for banteng population in Ujung Kulon National Park (UKNP). It is the largest population of banteng in Indonesia and is living in a limited habitat area. According to the best occupancy model, the most suitable areas for this species were the secondary forest located at the center portion of UKNP. The presence of the invasive cluster sugar palm, *Arenga obtusifolia*, in dry season provides additional alternative food for banteng when its main food is scarcer in the forest. Banteng was cathemeral all year round, with the proportion of cathemeral records and the recording rate did not change with the protection of the level area, moon phase or season. To reduce the probability of encountering predators, banteng avoided the space use of dholes. Selection and avoidance of habitats was stronger than avoidance of the predator activity areas. Habitat competition from domestic cattle which grazed illegally in the national park appears to be a problem to the species since zoonosis appears from domestic cattle to banteng. Therefore, effective law enforcement and an adequate conservation strategy are required to eliminate the impacts of both direct and indirect threats.

1. Introduction

Besides habitat destruction, intrinsic factors such as low reproduction rates, low densities, and extensive area requirements cause a high level of threat in the medium-large bodied mammal populations (Ripple *et al.* 2015; De la Torre *et al.* 2018). Approximately 60% of medium-large bodied ungulates are commonly faced dramatic population declines and habitat destruction and threatened with extinction (Ripple *et al.* 2015). Moreover, nine South East Asian ungulates species are listed as Critically Endangered (IUCN 2018). Several plants and wildlife species, mainly carnivores, are very dependent on the presence of ungulates species in nature. The

loss of ungulate species will cause tiered effects on ecosystems and in the future may affect essential economic and social services for humans (Ripple *et al.* 2015; Ripple *et al.* 2016). Amid a variety of threats to species, an increase in long-term research that addresses conservation solutions for these medium-large mammals are needed, particularly for lesser-known or non-charismatic species (Rahman *et al.* 2018).

The natural history and conservation need of many tropical forests of ungulates are poorly known, particularly in *Bos javanicus* (banteng). This lack of knowledge is heavily impacted by the conversion and degradation of lowland forests and illegal hunting, commercial trade, and bush-meat consumption (Nowak 1999; Di Marco *et al.* 2014). Banteng has its crucial function in ecosystems as dispersal vectors of plant propagules and nutrients (Bakker *et al.*

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2015), key modifiers and maintainers of ecosystem dynamics and habitat complexity (Jones *et al.* 1994; Waldram *et al.* 2008), as well as prey for medium to large carnivores (Mondal *et al.* 2012; Pudiyatmoko 2018; Rahman *et al.* 2018).

Banteng is a primitive ungulate with three subspecies and now are habituating the forests of Borneo, Java, and Bali in Indonesia. This species came from the mainland of Southeast Asia, particularly in Yunnan, Southern China through Peninsular Malaysia (Timmins *et al.* 2008). Today, banteng is extinct in South Malaysia, Bangladesh, and India (Medway 1969). Banteng is listing as an endangered species (the IUCN Red List; Gardner *et al.* 2016) and protected by Indonesian law (Peraturan Menteri Lingkungan Hidup dan Kehutanan Republik Indonesia No. P.106/MENLHK/SETJEN/KUM.1/12/2018; Indonesian Ministry of Environment and Forestry 2018) as Appendix I in CITES. Furthermore, Banteng is also listing as one of the 25 endangered species that need special attention to conservation (decree no. SK.180/IV-KKH/2015; Indonesian Ministry of Environment and Forestry 2015).

Here, we analysed a long-term study of 14-months intensive camera trap recording on the ecology and behaviour of banteng in the lowland tropical forest, Ujung Kulon National Park (UKNP). We aimed to: (1) provide relative encounter rates for recorded species, (2) investigate which habitat and environmental factors influence relative encounter rates, (3) observed the different time uses by banteng in their activity to avoid dangerous encounters with potential predators, and (4) investigate activity patterns, social structure and reproductive patterns.

2. Materials and Methods

2.1. Study Area

UKNP is a remnant of ancient forests that survived after a major eruption of a super volcano of Krakatau and tsunami in 1883 (Hommel 1987; Fauzi and Stoops 2004), and still have a significant threat of natural catastrophes such as earthquakes (Giachetti *et al.* 2012). It has a secondary growth tropical lowland rainforest with four main habitat types (Figure 1) and highly rainfall seasonal monsoonal climate. Biodiversity on land are suffering from anthropogenic activities, whereas the largest carnivores Javan tiger, *Panthera tigris sondaica*, has been extirpated, while the two remaining carnivores are Javan leopard (*Panthera pardus melas*) and dholes (*Cuon alpinus*) (Rahman *et al.* 2018). In UKNP, banteng (*Bos javanicus*) is one of six ungulate species and most prominent endemic mammals. The population of banteng are believed to be small in the UKNP (Pudiyatmoko 2005). In 1970, there were 13 locations known as banteng habitat in Java, whereas it shrinks into four locations which the

most abundant population occurs in UKNP recorded in 1997 (in total 905 individuals; Pudiyatmoko 2005).

2.2. Database Collection

We were superimposed 1/km² grid cell with total 329 grids at the study area and place cameras as close as possible at each intersection between the horizontal and vertical lines which comprised the grid (Figure 2). We used 108 camera-traps

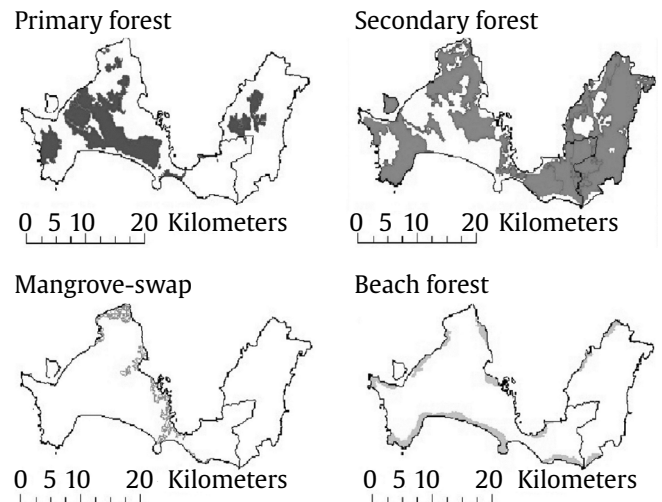


Figure 1. Four main habitat types in Ujung Kulon National Park

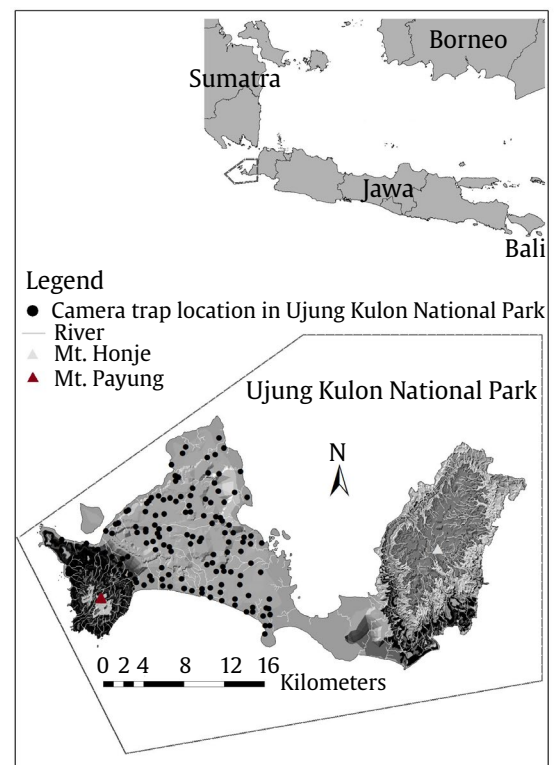


Figure 2. Camera trap locations in Ujung Kulon National Park (Java Island)

(Bushnell Trophy Cam 119467 and Bushnell Trophy Cam 119405) between December 2015 to January 2017 (with sampling effort=46,116 trap days) and were set to detect animals throughout the 24-hour/day, followed by c. sixty seconds of video, for each trigger. We set the time for every 10 seconds between successive triggers. We did not bait the camera, and all videos are with the information of date and time.

We randomly selected 134 sampling sites which designed represent all the four main vegetation types within UKNP. This scheme was designed to model the banteng's occupancy probability in the area UKNP, as well as documented its activity pattern. We were randomly drawn a grid over a digital map of the reserve and identified the sampling points by selecting its grid cells whereas at random one grid cell was one camera trap. This relatively even distribution of points within the UKNP maintained independence among points (a distance of <1 km). Cameras were set 170 cm above the ground with a 10-20 degree angle lead to the ground (following the standard design of camera trapping by the Rhino Monitoring Unit [RMU] team; Haryono *et al.* 2014). We did not consider any subsequent photograph of the same species which taken within 30 minutes interval as a new photographic event. Data were

managed using camtrap R software (Niedballa *et al.* 2016). Each photograph was located using latitude-longitude coordinate and converted into digital data in GIS using the ArcMap program.

2.3. Data Analysis

2.3.1. Relative Abundance Index

We report relative abundance for banteng encountered using Relative Abundance Index (RAI). We define all detections for a specific species summed for all camera traps over all days, multiply it with 100, and divide it by the total number of camera trap nights (O'Brien *et al.* 2003). Regarding un-normal distributed data or not transformed, we used nonparametric Kruskal Wallis tests and Mann-Whitney U post-hoc tests to check the differences in RAI between seasons.

2.3.2. Landscape Covariates and Occupancy Models

Fifteen variables were accounting for environmental traits, whereas four others for human impacts (Table 1). We tested the covariates for multicollinearity using Pearson's correlation matrix (STATS package R 3.1.1). We did not include covariates (with correlation >0.5) in the same candidate model.

Table 1. The environmental variables was considered to influence banteng distribution in Ujung Kulon National Park

Environmental variables classification	Variables	Measured analysis
Vegetation variables	Distance to primary, secondary forest edge, distance to invasive of cluster sugar palm <i>Arenga obtusifolia</i> (langkap), and vegetation productivity	Distance to the primary, secondary forest edge, and invasive of cluster sugar palm were extracted in ArcGIS 10.2.2, whilst vegetation productivity measured as the normalized difference vegetation index (NDVI, cf. Hansen <i>et al.</i> 2009)
Geomorphology variables	Elevation, slope, and distance to nearest river	The first two variables were computed on a 1 km ² grid resolution by averaging information extracted from a 90 x 90 m Digital Elevation Model (DEM; Landsat 8) (http://earthexplorer.usgs.gov ; http://srtm.csi.cgiar.org). Slopes were generated using slope function in ArcGIS (Jarvis <i>et al.</i> 2008). Distance to the nearest river was extracted in ArcGIS 10.2.2
Predation	Distance to the nearest presence of <i>C. alpinus</i> and <i>P. pardus melas</i>	Distance to the nearest presence of <i>C. alpinus</i> and <i>P. pardus melas</i> were extracted in ArcGIS 10.2.2
Climate variables	Annual rainfall, rainfall of the wettest month, rainfall of the driest month, annual mean temperature, the maximum temperature of the warmest month, and minimum temperature of the coldest month	These variables, derived from monthly temperature and rainfall values recorded between 1950 and 2000 from a global network of climate stations, were downloaded from the WorldClim database (http://worldclim.org/bioclim). All layers were projected into WGS 1984 Zone 48 South
Human impact variables	Distance to the nearest settlement, cultivated area, road, and illegal activity by human	Distance to the nearest settlement, cultivated area, road, and illegal activity by human (UKNP 2017) were extracted in ArcGIS 10.2.2

We classified camera-trapping data using 7-day intervals (26 occasions per season; Mackenzie *et al.* 2006). We estimated the site occupancy (Ψ) and detection probability (p) for banteng with three possible results: (1) the site was occupied, and the banteng was detected ($\Psi \times p$), (2) the banteng was present but not detected ($\Psi \times [1 - p]$), and (3) the banteng was not present and not detected ($1 - \Psi$).

We sampled 26 occasions from each site and estimated its detection probabilities. We used a maximum likelihood estimation of the proportion of sites occupied (Ψ) during the sample period as the projected parameter probabilities with a single-season model, respectively in wet and dry seasons. Because banteng shows a slow reproductive cycle (approximately a 295-day gestation period), we assumed zero population rate throughout the entire sampling period (Nowak 1999).

Habitat preference identification were constructed by a set of 214 candidate models with four different approaches: (1) occupancy probability and detectability as constant across all sites, (2) variation in occupancy as a function of covariates, (3) variation in detectability as a function of covariates, and (4) both variation in occupancy and detectability as a function of covariates. This model allowed us to evaluate the differences in habitat occupancy as determined by a single covariate or a set of covariates, which would contribute to an improvement in the model's performance.

We used PRESENCE 11.1 software (Mackenzie and Royle 2005) for occupancy modelling, and it was run using 2000 bootstraps to assess the adjustment fit (P) and the overdispersion parameter (\hat{c}). We also ranked the models by AIC adjusted for small sample size (AICc) following Burnham and Anderson (2002). All models with a $\Delta AICc$ value < 2 were considered to be competitive. We also used the AICc weight (w_i) for each model, which corresponds to the amount of evidence in favour of a given model, to choose the best model that we used to test our hypotheses.

2.3.3. Temporal Interactions Between Banteng-Potential Predator

The activity pattern of ungulates is strongly related to their predators (Dröge *et al.* 2017; Prugh *et al.* 2019). We followed Meredith and Ridout (2014) as a two-step procedure that resulted in a coefficient of overlap values from 0 (no overlap) to 1 (identical activity pattern). We used kernel

density estimation or by fitting a distribution from the flexible class of non-negative trigonometric sum distributions (Fernández-Durán 2004). For the second step, we calculated the overlap between two estimated distributions using the coefficient of overlapping, Δ (Ridout and Linkie 2009). Here, we applied the Δ_4 analysis because of the large sample size (Ridout and Linkie 2009; Meredith and Ridout 2014). We also used the overlap package (Meredith and Ridout 2014) to fit kernel density functions, estimate the coefficient of overlap, and calculate bootstrap estimates of the confidence intervals. We used Mardia-Watson-Wheeler Test (MWW test) to compare the distribution of detections across the diel cycle for all sampling campaign pairs, determining whether two activity patterns significantly differ (Batschelet 1981).

2.3.4. Daily Activity Patterns, Group Size and Group Patterns

Daily activity patterns and proportion of time spent active were estimated based on camera trap detections using Oriana circular statistics software (v4, Kovach Computing Services, Anglesey, UK) by applying the Rayleigh test (Batschelet 1981), the R package activity (Rowcliffe *et al.* 2014). We ran Wald tests for the statistical difference between activity level estimates at sunrise (dawn), sunset (dusk), noon and midnight. We used the Chi-square Test to see the frequency difference of observations between sunset-sunrise and between sunrise-sunset, with obtained timing from the Astronomical Applications Department of the US Naval Observatory (www.quickphase.com).

Furthermore, we used a moon calendar (Serrano 2011) to analyse the daily activity pattern on the lunar phase and continue with Analysis of Variance (ANOVA) to test the difference in activities rate between four moon phases (FM=full moon, FQ=first quarter, LQ=last quarter, and NM=new moon). We chose the two samples independent T-test to check whether there is a significant difference between activities on day and night as well as wet and dry seasons. To explain the distribution of activities in 24 hours, the effects of hours and night time (dummy variable where $D=1$ for 18.00-06.00, otherwise $D=0$) and to describe the interaction among activities, Generalized Linear Latent Variable Models (GLLVMs), a statistical model with random effects similar to multivariate Generalized Linear Mixed Models

(GLMMs), was used (Caraka *et al.* 2018). Warton *et al.* (2015) explained that the GLLVMs has several advantages in modelling multivariate-correlated responses compared to GLLMs; the number of parameters estimated is significantly smaller than that of GLLM due to the assumption imposed on GLLM to have an unstructured variance-covariance matrix. Besides, the latent variables used to incorporate correlation between responses can be used as a model-based approach to ordination. In this case, the responses are the number of Banteng activities including defecation, ingestive, locomotion, resting, social, and vigilant counted hourly (Table 2). Therefore, the multivariate responses are assumed to follow Poisson distribution. The mean responses and a linear predictor relationship are defined by:

$$E(Y_{ij}) = g(\mu_{ij}) = \eta_{ij} = \alpha_i + \beta_{0j} + \mathbf{x}'_i \boldsymbol{\beta}_j + \mathbf{u}'_i \boldsymbol{\lambda}_j$$

where $g(\cdot)$ was a link function corresponding to the selected distribution for responses, log link for the Poisson model. Latent variables, u'_i were assumed to have a standard normal distribution and independent of each other such that the variance-covariance matrix of random effects is $\Sigma = \lambda' \lambda$. The term α_i was the row effects representing the site variation and treated as fixed effects (Hui *et al.* 2015). We then used a `gllvm` package from R for fitting the model (Niku *et al.* 2017). For analysing group sizes and group patterns, we use descriptive statistics unless indicated differently.

3. Results

3.1. Relative Abundance Indices (RAI)

We documented 3,259 photographs of banteng with 817 and 644 independent photographs in the wet (RAI=3.10) and dry season (RAI=3.24), respectively. There are no significant differences between the RAI of wet and dry seasons for banteng ($W=3,427$; $p=0.367$).

3.2. Landscape Covariates and Occupancy Models

We observed the species in 96 (naive occupancy=0.71 and detectability=0.52) and 73 (naive occupancy=0.54 and detectability=0.41) of the 134 sites in wet and dry seasons, respectively. The center portion of UKNP resulted in the highest occupancy rates and detectability of bantengs ($\Psi=0.58-0.95$, $p=0.31-0.48$; $\Psi=0.44-0.88$, $p=0.18-0.48$; for wet and dry season, respectively). The number was followed by the southern portion of UKNP ($\Psi=0.32-0.43$, $p=0.18-0.29$; $\Psi=0.29-0.38$, $p=0.11-0.25$; for wet and dry season, respectively). The west portion of UKNP showed the lowest occupancy rates

Table 2. Ethogram for *Bos javanicus* activity states used in the study

Types of behaviour	Description
Excretion	Defecating or urinating
Locomotion	Moving between foraging source or within study area
Ingestive	Actively ingesting food or drink, or processing (chewing) food items during a grazing bout and or food searching with head below the vertebral column
Resting	Standing or sleeping in the sun or shade, neither ruminating nor scanning its environment
Socializing	Social interaction (necking), scratching, stroking, massaging self or others, nursing, and mating
Vigilant	Individuals scanning their surroundings and exhibiting agonistic displays

and detectability ($\Psi=0.08-0.13$, $p=0.06-0.09$; $\Psi=0.08-0.12$, $p=0.05-0.13$; for wet and dry season, respectively).

Our data best described the occupancy by two covariates in wet and dry seasons, respectively (Table 3). In wet season, distance to secondary forest edge had a negative relationship in which occupancy decreased as the distance to secondary forest edge increased ($\Psi=0.01-0.98$; Figure 3a), followed by distance to illegal activity which had a positive relationship, in which occupancy by bantengs increased as the distance to illegal activity by human increased ($\Psi=0.01-0.89$; Figure 3b). In dry season, distance to langkap had a negative relationship in which occupancy decreased as the distance to langkap increased ($\Psi=0.18-0.88$; Figure 3c), followed by distance to the nearest river which had a negative relationship, in which occupancy by bantengs decreased as the distance to the nearest river increased ($\Psi=0.10-0.96$; Figure 3d). Furthermore, detectability was affected by four covariates in both wet (Figure 4a) and dry seasons (Figure 4b).

3.3. Temporal Interactions Between Banteng-Potential Predator

The results of the MWW test revealed that patterns of daily activities were significantly different in all pairs of species (Table 4, Figure 5). The values of temporal overlap coefficients were <0.69 for all pairs of animals, which means that there was either low or intermediate overlap in their activity patterns. The most similar is between banteng and dhole, which are cathemeral species. Banteng and Javan leopard presented different peak activities. The banteng was most active in early morning and late evening, whereas the peak activity of Javan leopard was in midday.

Table 3. Occupancy models for Bantengs in the Ujung Kulon National Park, Indonesia, estimated by camera trapping between January and December 2016, grouped in sampling intervals of 7 consecutive days. With (.)=as constant, Ψ =occupancy, p =detectability, AICw=Akaike weight, and \hat{c} =dispersion parameter

Model	AICc	Δ AICc	AICw	Number of parameters
Wet				
Ψ (Secondary; illegalact); p (primary; illegalact; cuon; secondary)	568.90	0	0.30	8
Ψ (Secondary; illegalact); p (primary; cuon; secondary)	570.48	1.58	0.18	7
Ψ (Secondary; illegalact); p (primary; illegalact; cuon)	571.50	2.60	0.12	7
Ψ (Secondary; illegalact); p (primary; secondary)	571.63	2.73	0.09	6
Ψ (Secondary); p (primary; illegalact)	571.80	2.90	0.07	5
Ψ (Secondary; illegalact); p (cuon; secondary)	572.30	3.40	0.06	6
Ψ (Secondary); p (cuon; primary)	572.45	3.55	0.06	5
Ψ (Illegalact); p (cuon; primary)	574.70	5.80	0.05	5
Ψ (Illegalact); p (primary)	576.45	7.55	0.01	4
Ψ (Secondary); p (primary)	580.32	11.42	0.001	4
Ψ (.); p (.)	580.79	11.89	0.001	2
Ψ (.); p (cuon; primary)	581.40	12.50	<0.001	3
Ψ (Secondary); p (.)	581.84	12.94	<0.001	3
Ψ (Illegalact); p (.)	581.99	13.09	<0.001	3
Ψ (.); p (primary)	585.06	16.16	<0.001	3
Ψ (.); p (cuon)	586.70	17.80	<0.001	3
Ψ (Primary; elevation); p (primary)	587.18	18.28	<0.001	5
Ψ (Cuon); p (.)	587.60	18.70	<0.001	3
Ψ (Primary); p (.)	593.78	24.88	<0.001	3
Ψ (Elevation); p (.)	595.40	26.50	<0.001	3
Dry				
Ψ (Langkap; river; primary); p (primary; illegalact; cuon)	522.76	0	0.77	8
Ψ (Langkap; river; primary); p (primary; river)	523.81	1.05	0.41	7
Ψ (Langkap; river; primary); p (primary)	524.42	1.66	0.13	6
Ψ (Langkap; river); p (primary)	524.78	2.02	0.04	5
Ψ (Langkap; river); p (illegalact)	525.20	2.44	0.03	5
Ψ (Langkap); p (primary; illegalact; cuon)	527.13	4.37	0.006	6
Ψ (Langkap; elevation; primary); p (elevation; cuon; cultivated; illegalact)	528.74	5.98	0.005	9
Ψ (Primary); p (cuon)	531.30	8.54	0.003	4
Ψ (Cuon); p (.)	531.53	8.77	0.001	3
Ψ (Cuon; elevation; primary); p (primary; illegalact)	532.98	10.22	<0.001	7
Ψ (.); p (.)	534.24	11.48	<0.001	2
Ψ (Cultivated); p (.)	535.76	13.00	<0.001	3
Ψ (Elevation); p (.)	536.63	13.87	<0.001	3
Ψ (Elevation; langkap); p (cultivated)	538.78	16.02	<0.001	5
Ψ (Illegalact; cultivated); p (.)	541.01	18.25	<0.001	4
Ψ (.); p (primary)	541.76	19.00	<0.001	3
Ψ (.); p (cuon)	543.08	20.32	<0.001	3
Ψ (Primary; cultivated); p (primary; illegalact)	544.30	21.54	<0.001	6
Ψ (Cuon); p (.)	555.16	32.40	<0.001	3
Ψ (Primary); p (cultivated)	564.30	41.54	<0.001	4

3.4. Daily Activity, Group Sizes and Group Patterns

The number of activities is higher during the dry season compare to wet season, although there is no significant difference in the activity rate between these two seasons ($t=-1.988$, $df=46$, p -value=0.052). Numbers of banteng's records in wet and dry seasons ($n=817$ and 644 , respectively) indicated a similar overall activity level in each season (0.72 and 0.68, respectively). Also, there was no significant deviation has been detected from a uniform activity distribution throughout the day in both wet and dry seasons ($z=3.13$, $p=0.101$ (Figure 6a); $z=5.43$, $p=0.455$ (Figure 6b), respectively) using the Rayleigh tests, suggesting

that this species may be cathemeral. Nevertheless, banteng tended to be more active at dusk than dawn ($p=0.012$), with a mean activity time of $16:42 \pm 18:48$ circular stdev. There was no significant difference in number of encounters between day and night time in both wet and dry seasons ($\chi^2=2.43$, $df=1$, $p=0.119$; and $\chi^2=3.38$, $df=1$, $p=0.227$, respectively).

The number of activities occurred at four moon phases shows indistinct differences from one to another, where the highest activity is between 17.00 and 18.00. This finding is also supported by the statistical test showing that there is no significant difference in activities between four moon phases

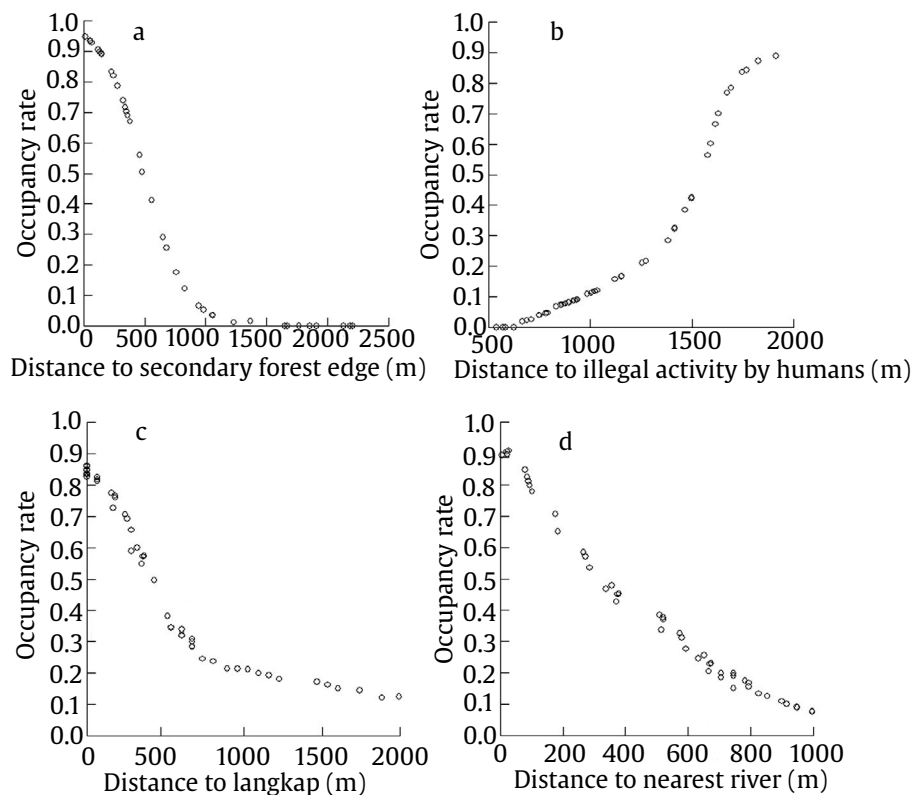


Figure 3. Relationship between occupancy of the tropical lowland forest, Ujung Kulon National Park, Banten, Indonesia, by *Bos javanicus* and distance to secondary forest edge (a) and distance to the nearest illegal activity by human (b) in wet season. Relationship between occupancy of the tropical lowland forest, Ujung Kulon National Park, Banten, Indonesia, by *Bos javanicus* and distance to invasive of palm species: Langkap *Arenga obtusifolia* (c) and distance to the nearest river (d) in dry season

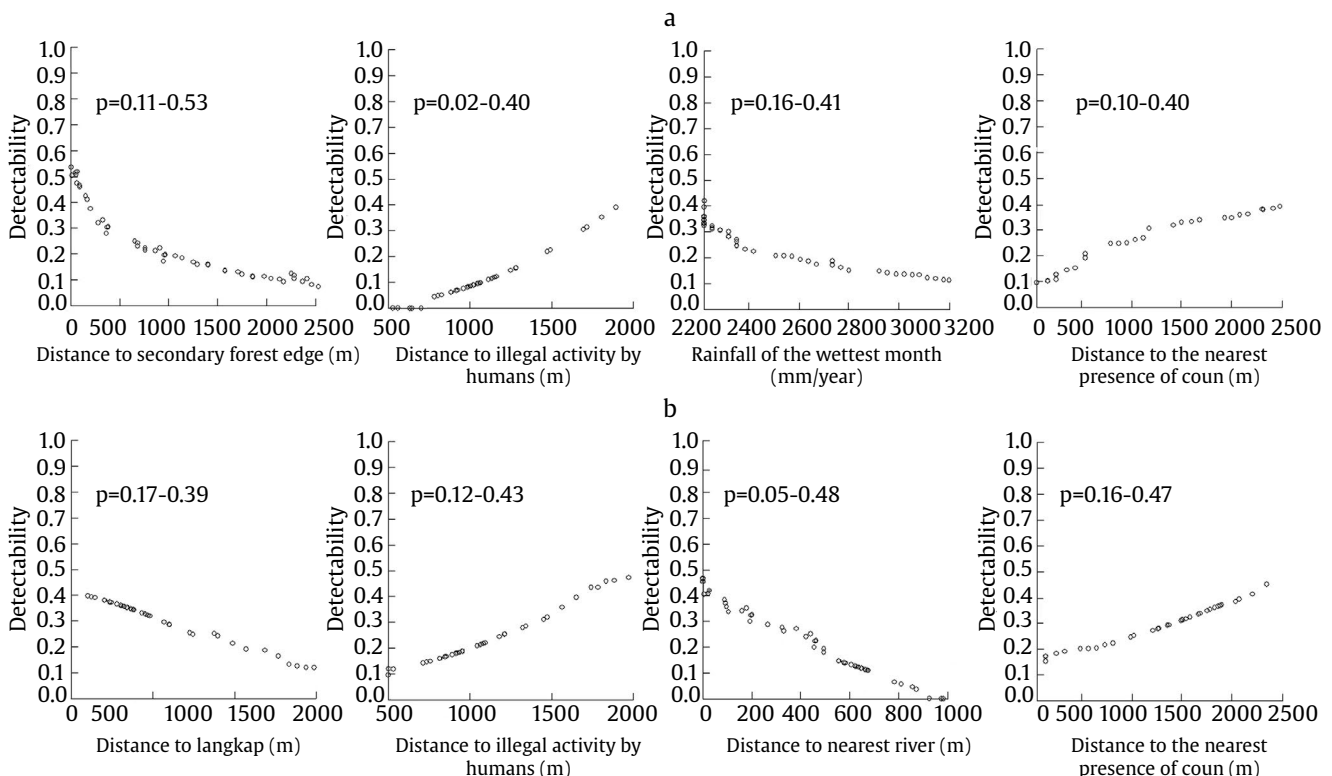


Figure 4. Relationship between detectability of *Bos javanicus* Ujung Kulon National Park, Banten, Indonesia (a) in wet season, and (b) dry season

Table 4. Overlap coefficient (Δ_4) between prey-predator activity patterns, 95% confidence intervals (95% Cis), and Mardia-Watson-Wheeler (MWW) test

	Δ_4 (95% ci)	MWW	p value
Banteng \times Javan leopard	0.56 (0.53-0.59)	39.45	<0.0001**
Banteng \times Dholes	0.69 (0.66-0.72)	8.30	<0.05*

The p value of less than 0.05 indicate that the two sets of circular distributions come from different distribution

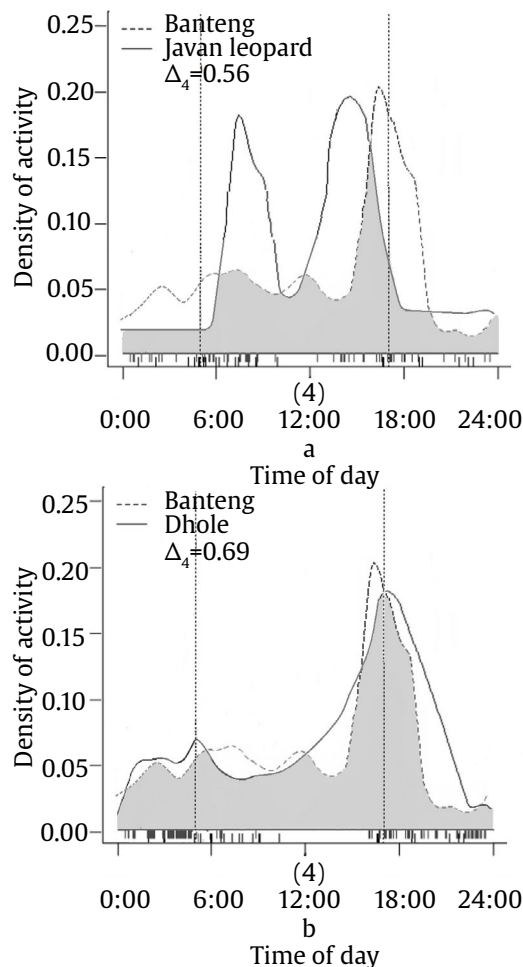


Figure 5. Daily activity patterns and overlap of (a) banteng-Javan leopard, and (b) banteng-Dhole

($F=0.328$; $df\ 1=3$, $df\ 2=92$, $p\text{-value}=0.805$, Figure 7a). On full moon phase (FM), there is no meaningful increase in activities during the night which is also supported by the statistical test ($t=1.073$, $d.f.=1$, $p\text{-value} = 0.295$, Figure 7b).

Our finding shows that activities such as excretion, locomotion, ingestive and vigilant are found to be positively-strongly correlated indicating these activities tend to happen at the same times. Some activities such as excretion, resting and vigilant, however, show a negative association with social

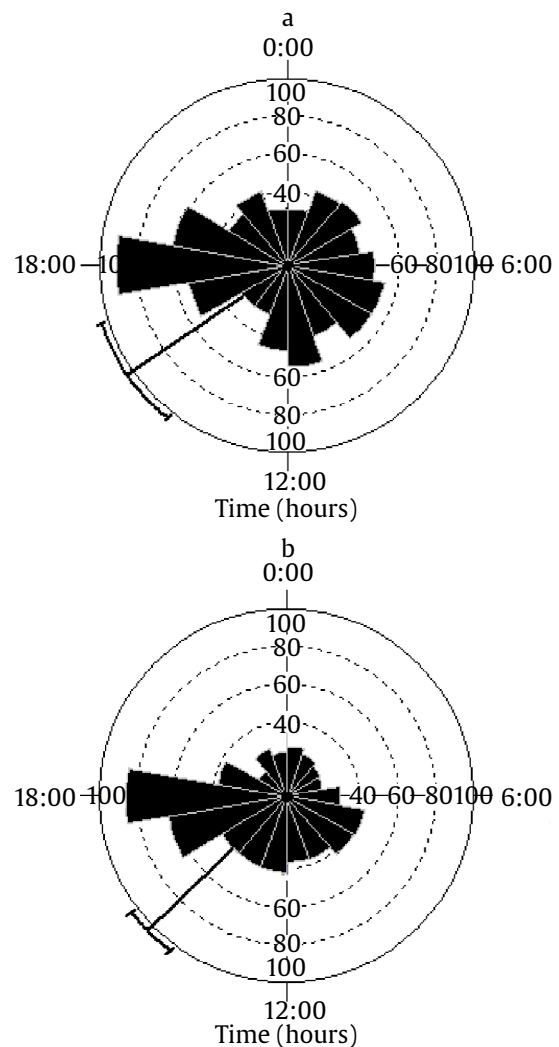


Figure 6. Activity plots based on camera trap records of *Bos javanicus* (a) in wet season and (b) in dry season

activity explaining that they occur in the different time with social activity. We found two points of time where the activities rate reaches its peak (between 17.00-19.00). In the night time variable, we found only ingestive activity receives significant effects where it indicates the species tend to do ingestion during the day, the sign of parameter is negative. The correlogram matrix displays the interaction between activities for 24 hours (Figure 8).

Overall, bantengs appeared in small family groups (with or without males), pairs, or as a single male. Changes in group structure over the year indicate a peak of the reproductive stage in the wet season, particularly in January (Figure 9). The mean group size, mean litter size, and group combination are shown in Table 5.

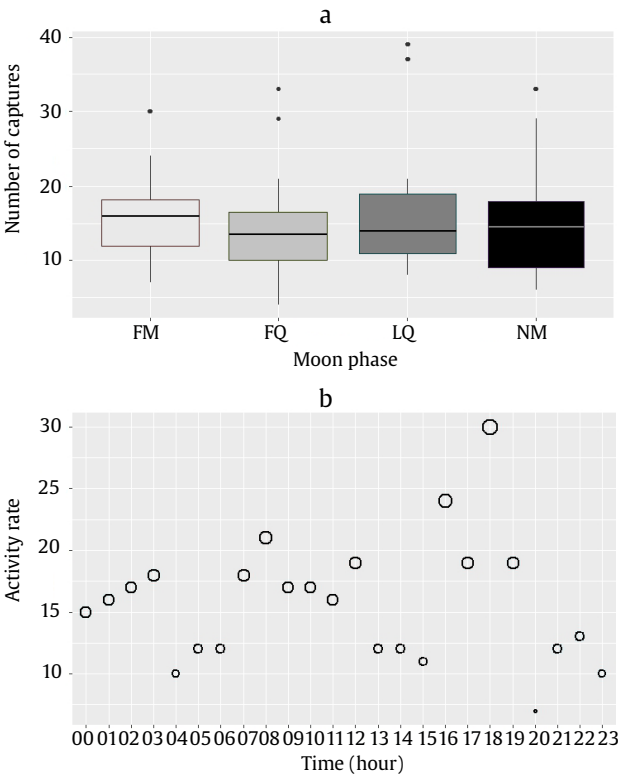


Figure 7. The number of *Bos javanicus* activities occurred at four moon phases (a) and the number of activities on full moon phase within 24 hours during a year (b)

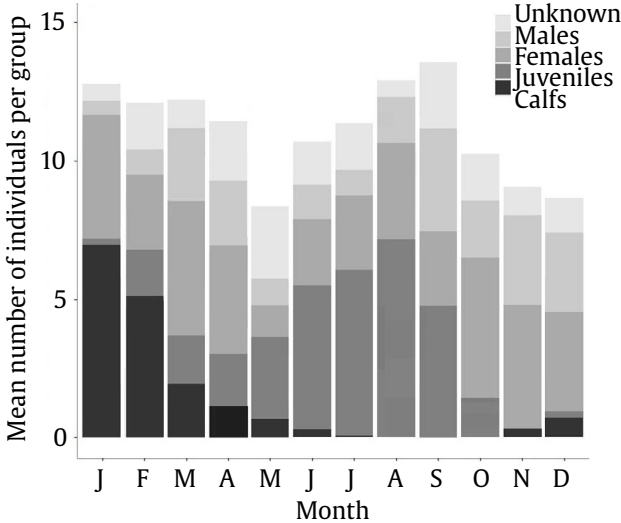


Figure 9. Annual patterns in group size and structure based on camera trap records of *Bos javanicus*

Table 5. Mean group size, mean litter size and group combinations of banteng (*Bos javanicus*) recorded in 30-second videos in December 2015-January 2017

Parameters	
N of videos	1461
Mean group size±sd (range)	7.54±6.50 (1-15)
Litter size±sd (range)	3.38±2.14 (1-6) (n=269)
Number of adult females (range)	2.65±1.98 (0-7)
Number of adult males (range)	1.18±0.44 (0-3)
Number of unknown adults (range)	2.10±1.04 (0-6)

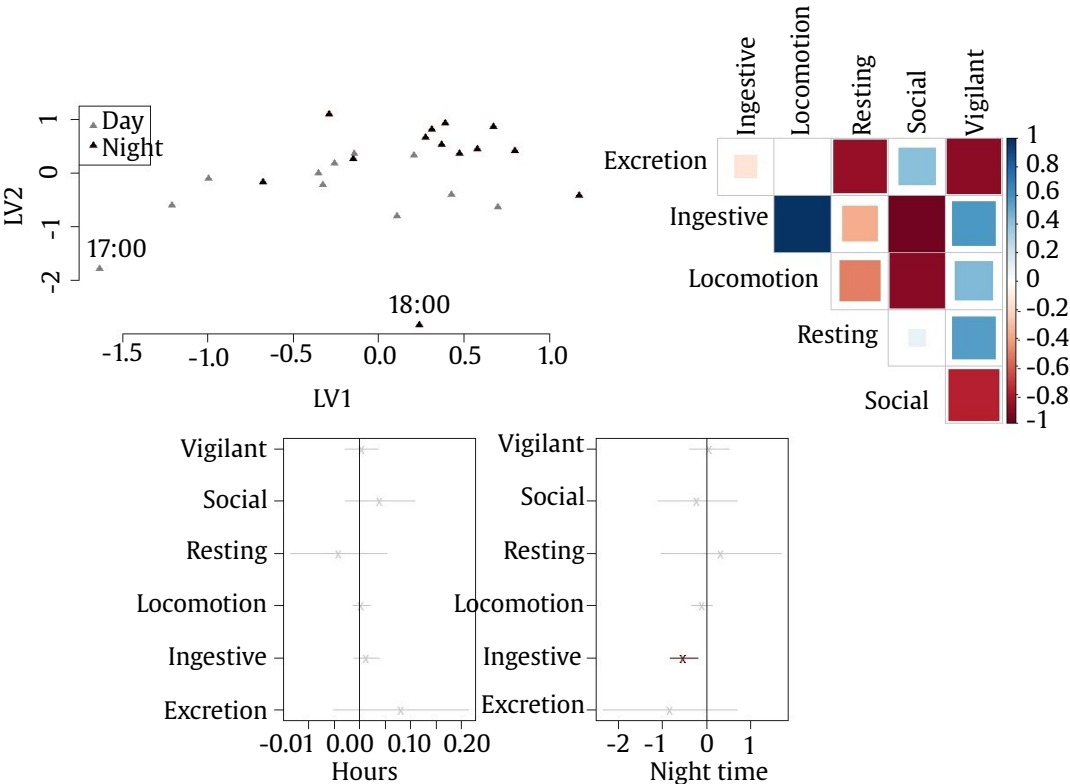


Figure 8. The distribution of total kind of activities in 24 hours during 2016 by *Bos javanicus*

4. Discussion

Although our data covers only a single national park on the western tip of Java Island, the robust sampling design and the variety of surveyed habitats support a more considerable extent to our results. This may constitute a basic knowledge for other areas in Java and the whole species' range.

We detected banteng almost at all sites in the photographs, including sites where no banteng has been reported before. Using occupancy models, we have been identifying suitable habitat for bantengs to establish conservation priorities for the species in the UKNP. Through this study, we proposed the conservation priority of banteng's habitat that integrates occupancy models and landscape information at local scale for the first time. Identifying areas in wet and dry seasons with high presences of banteng population will be a valuable information tool to determine conservation priority areas in UKNP. It includes subsequently direct ranger patrols, anti-poaching efforts, and anti-encroachment operations. The suitability distribution areas generated by our models show occupancy areas of the high and low presence for banteng. Also, it provides an understanding of relevant natural and anthropogenic variables which support the occurrence of the banteng.

Our results showed the landscape and anthropogenic variables which best describe the occupancy of banteng in the UKNP. These results supported our predictions that banteng occupancy in UKNP was positively associated with the primary source of food and negatively with disturbance by humans. Our result showed a lower number of records in the wet season compared to the dry season, which possibly related to a lower level of activity caused by higher food availability in the secondary forest. Thus, the food and water source availability during the dry season is scarcer. This relative scarcity is leading to a broader movement of banteng to the rivers and langkap areas which increasing the encounter probability (Buchholz 1990; Rahman *et al.* 2017).

Several studies carried on in other regions have reported the association between banteng and its areas near water (Manh-Ha 2009). Thus, the highest occupancy rates and detectability of the banteng in the center portion of the UKNP attributed to the higher quantity of water portion in the park compared to the other portions of UKNP (i.e., south and west). Moreover, the distance to water was one of the covariates that best described the distribution of bantengs (Imron *et al.* 2016; Chaiyarat *et al.* 2018). Currently, UKNP is experiencing a lower water level during the dry season

because lack of rainfall, they will use water sources such as rivers that still have water or use permanent wallows that are usually used by rhinos. Additionally, mineral licks are also an essential feature of banteng's habitat. At the coastal regions which lack minerals, animals licking behaviour will meet the mineral requirements by occasionally drinking seawater (Halder 1976).

Hommel (1987) recorded no palms *Arenga obtusifolia* (langkap) in UKNP in the 1980s, but it is abundant along the area nowadays (Hariyadi *et al.* 2012). Langkap is an invasive plants species and has grown dominantly in the area where the originals banteng food source occurred. We suspected langkap as an alternative food source for banteng. Previous studies have indicated langkap dominancy as a threat to rhinos' natural food availability (Nardelli 2016). Thus, occupancy and detectability models demonstrate high occurrence probabilities in such habitat areas.

UKNP is a national protected area (Ministry of Forestry 1992) and is the UNESCO World Heritage Site since 1991, but illegal pastures for livestock, forest encroachment, and poaching are still common practices in the area (UKNP 2017). As much as 299 cases of illegal activity occurred in the wet season (UKNP 2017). Other studies carried out at Baluran National Park (and also Meru Betiri NP and Alas Purwo NP) in East Java have shown that occupancy and abundance of this banteng are negatively correlated with the level of poaching (Pudyatmoko 2004). Based on the interview that we conducted with local people, they are ignoring the bantengs. Here, bantengs are not the hunting target because they only cause limited damage to crop in the surrounding area. The biggest threat for bantengs in UKNP is illegal livestock grazing activities which lead to zoonoses (Radcliffe *et al.* 2016; Khairani *et al.* 2018).

Besides illegal activity by humans, the distance to forested patches, rainfall and predation may also influence the detectability of banteng in UKNP. The high degree of rainfall and predation by dhole in the lowlands could be associated with the lowest rates of occupancy and detectability of bantengs (Rahman *et al.* 2018). Furthermore, the detectability of bantengs increased as the distance to deforested patches decreased, which indicated that bantengs accosted the sites nearest to deforested patches. In UKNP, dhole has a significant positive interaction and overlaps with the banteng, whereas banteng is captured as prey by dhole (Rahman *et al.* 2018). Dholes have the advantage of large-sized prey hunts because they are living in groups. We need to consider on the spatiotemporal interaction between dholes and banteng. This relationship was the only pair of species interaction which represented a strong

negative spatial interaction and temporal segregation. Bantengs in the Baluran National Park is the only large mammal that exhibits significant negative spatial interaction with free-range livestock (Pudyatmoko 2017). Furthermore, dhole might be responsible for population declines of banteng in Alas Purwo National Park (Hedges and Tyson unpublished work). This finding indicates the banteng sensitivity to predation. Dholes' exhibit similar predator to prey ratios, in contrast to solitary hunters' Javan leopard (feed on the equal or slightly smaller size) (Hayward *et al.* 2014; Rahman *et al.* 2018). The pattern of movement and distribution of herbivore animals is in general strongly associated with predators. They tend to adjust the movement through the avoidance strategy (Flaxman *et al.* 2011; Rosier and Langkilde 2011).

Here, we found that bantengs are active both at night and during the day, which in line with the previous study in UKNP (Halder 1976; Timmins *et al.* 2008). Moreover, banteng in UKNP tended to be more active at dusk than dawn (mean activity time between 17:00-19:00). In undisturbed areas, banteng does not differentiate much in their activities pattern between day and night (Hoogerwerf 1970), which we can see in Baluran National Park and Alas Purwo National Park (Pudyatmoko 2005). Both in Baluran and Alas Purwo National Park, several groups graze in the open area at dusk until 08:00, and occasionally until 10:00. Afterwards, banteng will enter the forest to avoid the hot temperatures in the open area. They will come back to the open area in the afternoon (around 15:00) and stay until the evening. In contrast with the two locations above, differential use of habitat between day and night has not yet been documented for banteng in UKNP. Here, the individuals are resting and foraging in dense vegetation throughout the day. Only if in some periods the food is abundant in the open area (such as pasture), banteng will utilise this location. Our finding shows that excretion, locomotion, ingestive, resting, socialising, and vigilant are roughly identical during both the day and the night in 24 hours in a year.

Since our results showed bantengs activities within 24 hours a year with no differences, it also showed no correlation between moon phase to bantengs activities. Bantengs became adaptable to be active in the evening, but they are not a nocturnal species. It is different than some nocturnal ungulates species which significantly increase the frequency of foraging during a particular moon phase (Pratas-Santiago *et al.* 2016; Traill *et al.* 2016). Banteng is known as ruminants that carry out

foraging activities throughout the day. Ruminants have a unique digestive system which designed to ferment the digested food and provide precursors for energy, unlike monogastrics species such as swine and poultry (Orr *et al.* 2001).

Group composition of bantengs seems to be variable, with males sometimes staying in family groups when immatures are present. An old banteng usually leads each herd. Young bantengs form a herd with two or three individuals. Adult bantengs are solitary but enter a banteng herd for mating (Lekagul and McNeely 1988). Our mean group size of 3 (range 2-15) is significantly lower than the previously reported mean group size of 4 (2-30) (Lekagul and McNeely 1988; Gardner *et al.* 2014). The differences in the number of groups are possible due to differences in study areas. Most of the reported groups of bantengs are those that inhabit open areas such as pastures. In the open area, the banteng group appears to be a larger group size since it may be a combination of several small groups together in the open area. Supported by our study which explaining in detail the behaviour of banteng in open areas is to reduce threats by predators. Preys that live in larger groups can intimidate a predator by simultaneously mobbing it, driving the predators away before it can attack. Prey can also behave by signalling the predator, that it would be a waste of time and energy attempting an attack (Kie 1999; Rosier and Langkilde 2011).

Moreover, our data support the peak of reproduction phase which is showing the highest peak in the wet season (January and February), with no immatures observed during the end of the wet season (April and May). Increased reproduction in the peak of the wet season has been recorded from Cambodians living within Banteng habitat, which suggested that the majority of births occurred in December and January (S. Hedges unpubl data 1991-2002).

5. Conclusions

Ultimately, we propose the findings of our study to facilitate more effective and reliable occupancy monitoring of banteng in UKNP. Accurate patch-specific occupancy data will enhance our knowledge of the species distribution, enable tracking changes in population status, assist in the selection of additional management focal areas, and facilitate monitoring responses to habitat management. Therefore, we recommend our survey protocol and recommendations to be incorporated into an adaptive management

program for the species. Our findings also provide insight into the role of detection in monitoring rare and cryptic banteng that occupy dense habitats. Our approach may be useful in developing monitoring programs for conservation concern of other species, for which accurate patch-level data on occupancy status are necessary for conservation management. Moreover, further research is required to provide information for long-term conservation planning such as 'Project Banteng', akin to 'Project Rhino' and 'Project Elephant', for the future survival of the species.

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